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Description

This invention relates to the purification of propylene oxide. More particularly, this invention relates, to a distillation process for removing contaminating quantities of impurities including oxygen-containing impurities such as methanol, acetone and water from an impure propylene oxide feedstock. Still more particularly, this invention relates to a method wherein an impure propylene oxide feedstock contaminated with from about 50 to about 4,000 ppm of methanol, from about 0.01 to about 2 wt.% of water and from about 0.01 to about 2 wt.% of acetone is purified in an extractive distillation column using triethylene glycol as an extractive distillation agent.

It is known to react a hydroperoxide feedstock such as tertiary butyl hydroperoxide with propylene in the presence of an epoxidation catalyst in order to provide a reaction product comprising propylene oxide, an alcohol corresponding to the hydroperoxide feedstock, a solvent, and impurities (see, for example, Kollar U. S. Patent No. 3,350,422, Kollar U. S. Patent No. 3,351,635 and Sorgenti U. S. Patent No. 3,666,777.

It is also known to separate the reaction product by distillation in order to obtain a plurality of fractions including, for example, a propylene recycle fraction, a propylene oxide product fraction and an alcohol fraction.

It is also known that methanol, acetone and water are common contaminants for propylene oxide which are removed only with difficulty.

For example, Mitchell et al. U. S. Patent No. 2,550,847 is directed to a process for separating purified propylene oxide from a crude propylene oxide product contaminated with acetaldehyde, methyl formate, methanol, etc., by treating the crude mixture with an aqueous basic substance followed by recovery of the purified propylene oxide by any suitable means such as by decantation. Mitchell et al. reported a recovery of a product containing 78 to 82 wt.% of propylene oxide which, they stated, could be increased in purity to about 95 to 99% by fractional distillation.

Robeson et al. U. S. Patent No. 2,622,060 discloses a process for the purification of propylene oxide contaminated with impurities, including methanol, by subjecting the impure propylene oxide to distillation in the presence of an extractive distillation agent comprising an aqueous solution of an alkali. The inventors report in Example 1 of their patent a method wherein 500 parts by weight of a crude propylene oxide fraction was extractively distilled in accordance with their invention to obtain 325 parts by weight of a product containing about 99.6 wt.% of propylene oxide. Thus, a significant loss of propylene oxide occurred during the process.

In a process unrelated to the purification of propylene oxide, Goddin et al. in U. S. Patent No. 2,751,337 disclose a process for separating acetone from a mixture of acetone with methanol and methyl acetate utilizing water as an extractive distillation agent.

Hamlin et al. in U. S. Patent No. 3,409,513 disclose the hydro-extractive distillation of mixtures comprising acetone, lower aliphatic alcohols and esters of lower aliphatic alcohols with carboxylic acids. It is pointed out by the patentees that acetone, methyl acetate and methanol form an azeotrope boiling at 55.5-56.5 °C. Williams et al. propose to recover partially purified acetone from such a terniary azeotrope by liquid-liquid extraction with water followed by hydro-extractive distillation of the aqueous phase in order to obtain a partially purified acetone fraction.

Hoory and Newman U. S. Patent No. 3,632,482 is directed to a propylene oxide recovery process by extractive distillation using an alcohol-ketone-hydrocarbon solvent. The invention relates to a method for the purification of crude propylene oxide contained in a mixture produced by the epoxidation of propylene with an organic hydroperoxide and calls for extractive distillation of the crude propylene oxide in a plurality of successive extractive distillation zones with the aid of a solvent mixture consisting essentially of hydrocarbons, alcohols, and/or ketones corresponding to the organic hydroperoxide employed in producing the propylene oxide. In the preferred embodiment of their invention, the extractive distillation agent is a recycle fraction from a three column distillation sequence wherein the bottoms from the third distillation column are flashed to obtain an overhead composed of hydrocarbons, alcohols and/or ketones which is recycled as an extractive distillation agent to the three distillation columns involved in the propylene oxide purification sequence.

Burns et al. U. S. Patent No. 3,715,284 discloses a process for the purification of impure propylene oxide using acetone or a mixture of acetone with methanol which is introduced into a distillation column either below or together with the impure propylene oxide.

Schmidt U. S. Patent No. 3,881,996 is directed to a distillation sequence employing at least three and optionally four columns for the purification of crude propylene oxide, one of the columns optionally being an extractive distillation column wherein a hydrocarbon such as octane is used as the extractive distillation agent.

Schmidt U. S. Patent No. 4,140,588 discloses a process for the purification of propylene oxide contaminated with methanol and acetone using water as an extractive distillation agent, the water being introduced into the distillation column above the point of introduction of the crude propylene oxide feed.

Schmidt states at column 2, lines 50-55 that: "Propylene oxide, however, has a substantial solubility in water and is readily hydrolyzed to propylene glycol (PG) in the presence of large amounts of water"--i.e., in the reboiler section of the tower.

- U. S. Patent No. 4,971,661 discloses the use of an aqueous acetone extraction to remove methanol from propylene oxide.
- U. S. Patent No. 3,578,568 discloses the use of glycols or glycol ethers in an extractive distillation to remove oxygen containing impurities such as acetone, acetaldehyde, and methanol. It is claimed that the concentration of the solvent in the vapor space in the extractive distillation zone of the distillation tower is preferably between 15 and 50 mole percent of the total vapor.

Compared to U. S. Patent No. 3,578,568, this invention uses considerably lower concentrations of solvent in the extractive distillation zone to remove water and oxygen-containing impurities such as acetone. Since the concentration of the triethylene glycol is lower, the size and heat requirements of the associated triethylene glycol regenerator are reduced.

U. S. Patent No. 3,607,669 discloses the use of a C₈ to C₁₂ hydrocarbon to separate propylene oxide from water.

Shih et al. U. S. Patent No. 5,000,825 discloses the purification of monoepoxides such as propylene oxide that are contaminated with oxygenated impurities such as water, low molecular weight alcohols, low molecular weight ketones, low molecular weigh aldehydes and the like by the extractive distillation of the contaminated monoepoxide using a lower glycol containing 2 to 4 carbon atoms. Examples of lower glycols that are given in the patent include ethylene glycol, 1,2-propane diol, 1,3-propane diol, 1,4-butane diol, 1,2-butane diol, 1,3-butane diol and 2,3-butane diol. It is stated that higher diols or higher glycol ethers do not provide sufficient selectivity for the removal of such impurities and are not included as the extractive distillation solvents suitable for use in the invention.

An object of the present invention is to provide a distillation process for removing contaminating quantities of impurities including oxygen-containing impurities such as methanol, acetone and water from an impure propylene oxide feedstock.

The present invention provides an extractive distillation process for the removal of oxygenated contaminants, including water, methanol, and acetone from an impure propylene oxide feedstock comprising propylene oxide contaminated with water, methanol, and acetone, characterised by introducing the said impure propylene oxide feedstock into the lower half of a distillation column, introducing an extractive distillation agent comprising triethylene glycol above the impure propylene oxide feed point, at a ratio of feedstock to extractive distillation agent of from 1:1 to 20:1, withdrawing from said distillation column an overhead distillate fraction comprising of essentially anhydrous propylene oxide contaminated with reduced quantities of acetone and methanol, and a bottoms distillation fraction containing substantially all of the triethylene glycol, water and acetone and a portion of the methanol introduced into said distillation column. Preferably the impure propylene oxide is charged to the lower half of a distillation column containing at least 25 theoretical plates and the extractive distillation agent is added at a point at least 4 theoretical plates above the impure propylene oxide charge point. It is also preferred that the distillation conditions comprise a pressure of 69 to 276 kPa (10 to 40 psia), a reflux ratio of from 1:1 to 5:1, and a reboiler temperature within the range of 100 ° to 250 °C. and a top temperature of 20 ° to 80 °C.

When propylene is reacted in liquid phase with an organic hydroperoxide such as tertiary butyl hydroperoxide in solution in a solvent such as tertiary butyl alcohol in the presence of a soluble epoxidation catalyst such as a molybdenum epoxidation catalyst, a reaction mixture is formed comprising propylene oxide, an alcohol corresponding to the organic hydroperoxide feedstock and impurities including water and other oxygenated impurities such as methyl formate, acetaldehyde, acetone and methanol.

Propylene oxide is a hygroscopic substance, so that water is removed only with difficulty. It is important to remove as much of the water as possible, however, because the water present in the propylene oxide will tend to react with the propylene oxide to form propylene glycol.

It is also important to reduce the level of other oxygenated contaminants such as methanol and acetone to the lowest reasonably attainable level.

In accordance with conventional practice, an epoxidation reaction product formed by the molybdenum-catalyzed reaction of propylene oxide with tertiary butyl hydroperoxide in solution in tertiary butyl alcohol is separated into the principle components by distillation so as to form distillation fractions including a propylene distillation fraction, a propylene oxide distillation fraction, a tertiary butyl alcohol distillation fraction and a heavy distillation fraction containing the molybdenum catalyst and other products and by-

products of the epoxidation reaction. However, the distillation fractions that are thus-obtained are characterized by the inclusion of impurities and, normally, must be further treated if commercially acceptable products are to be obtained. This is especially true for a propylene oxide distillation fraction contaminated with water and oxygenated contaminants including methanol and acetone.

It has been surprisingly discovered in accordance with the present invention that substantially all of the water initially present in a contaminated propylene oxide feedstock can be removed therefrom when the propylene oxide feedstock is extractively distilled in the presence of an extractive distillation agent consisting essentially of triethylene glycol. Even more surprising is our discovery that substantially all of the acetone and most of the methanol present in the contaminated feedstock can also be removed from the propylene oxide when using triethylene glycol as the extractive distillation agent.

The invention will be further described with reference to the accompanying drawings. The drawing is a schematic flow sheet with conventional parts omitted showing the general recovery sequence that is used in accordance with the present invention in purifying propylene oxide.

In the drawing, for convenience, the present invention is illustrated in connection with a process wherein the propylene oxide is prepared by the epoxidation of propylene with tertiary butyl hydroperoxide in solution in tertiary butyl alcohol to provide a reaction product comprising propylene oxide and additional tertiary butyl alcohol.

Turning now to the drawing, there is shown a schematic flow sheet illustrating a preferred method of practicing the process of the present invention. In the drawing, conventional parts such as valves, pumps, temperature sensors, pressure sensors, heaters, coolers and flow control regulation apparatus have been omitted.

In accordance with a preferred embodiment of the present invention, propylene oxide is separated in a preliminary distillation zone (not shown) from other components of an epoxidation reaction mixture in order to provide an impure propylene oxide fraction contaminated with oxygen-containing impurities such as acetone, methanol and water.

The impure propylene oxide feedstock that is thus obtained in the preliminary distillation zone is then purified in a propylene oxide purification distillation zone, which in accordance with the preferred embodiment of the present invention, comprises two distillation columns, each of which is equipped with an appropriate reflux condensing means and an appropriate reboiler heating means.

In accordance with the present invention, an impure propylene oxide fraction contaminated with from about 50 to 4,000 ppm of methanol, from about 0.01 to 2 wt.% of acetone and about 0.01 to 2 wt.% of water and other oxygen-containing impurities is charged by way of a line 120 leading to a distillation column 100 which, in accordance with the present invention, will preferably be a column containing at least about 10 theoretical plates and more preferably, from about 30 to about 100 theoretical plates. The column 100 is suitably operated under distillation conditions including a pressure of about 69 to 276 kPa (10 to 40 psia), a reflux ratio of from 2:1 to 10:1, a reboiler temperature within the range of about 100 ° to 250 °C (e.g. 210 °C) and a top temperature of about 20 ° to 80 °C (e.g. 20 °C).

The impure propylene oxide is preferably charged to the distillation column 100 in the lower half thereof. An extractive distillation agent consisting essentially of triethylene glycol is charged to the upper half of the distillation column 100 by an extractive distillation charge line 106.

Essentially anhydrous purified propylene oxide containing about 100 ppm or less of water is removed from the column 100 as a light distillation fraction 112, the purified propylene oxide in the line 112 containing significantly reduced amounts of methanol and acetone, such as about 15 to 900 ppm of methanol and about 0.1 to 100 ppm of acetone. A heavier fraction 110 is withdrawn from the distillation column 100 which contains substantially all of the extractive distillation agent charged by the line 106 and also substantially all of the water, acetone and other oxygen-containing impurities introduced into the column 100 with the impure propylene oxide 120.

The heavier distillation fraction 110 from the column 100 comprising water, methanol, acetone, tertiary butyl alcohol and other impurities and extractive distillation agent is charged to a second distillation column 200 wherein light impurities such as methanol, acetone, water, etc., are separated overhead as a distillation fraction 204 that is discharged from the system for any suitable use, such as for use as a steam boiler feedstock or for recovery.

A heavier distillation fraction 106 is discharged from the distillation column 200 comprising triethylene glycol which is recycled to distillation column 100 by line 106.

Triethylene glycol is a compound having the formula:

HO-CH2 CH2-O-CH2 CH2-O-CH2 CH2-OH

Triethylene glycol is a colorless, hygroscopic, practically odorless liquid having a boiling point of 287.4 °C., a vapor pressure of less than 0.01 mm (20 °C.) and a specific gravity of 1.1254 (20/20 °C.). Technical grades of triethylene glycol will contain an appreciable amount of water (e.g., about 0.01 to about 0.1 wt.%). Therefore, if fresh technical grade triethylene glycol were introduced directly into the column 100, a substantial amount of undesired contaminating water would also be introduced. In accordance with the present invention, fresh triethylene glycol, either as the original charge, or as make-up solvent, is introduced into the system by a branch line 230 leading to the charge line 110 for the second distillation column 200 so that any water introduced into the system with the fresh triethylene glycol will be separated therefrom in the column 200 and withdrawn from the column 200 through the line 204.

Water and oxygen-containing impurities such as acetone and methanol are difficult to remove from propylene oxide by standard distillation. The use of extractive distillation columns with triethylene glycol as the solvent improves the separation of these impurities from propylene oxide.

EXAMPLES

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The invention will be further illustrated by the following non-limiting examples.

Example I

20 Comparative distillation runs without the use of an extractive distillation solvent and with triethylene glycol as the solvent are tabulated below. The runs were conducted in a 25 plate 2.54cm (1") Oldershaw type glass distillation column operating at atmospheric pressure.

Component	Feed wt%	Solvent wt%	Overhead wt%	Bottoms wt%
Water	0.1800	N/A	0.0784	01.3900
(WATER FREE BASIS)				
Lighter than MeOH/MF	ND	N/A	ND	N/An
MeOH/MF	0.207	N/A	0.119	N/An
PO	99.184	N/A	99.881	N/An
Acetone	0.524	N/A	ND	N/An
TBA	0.084	N/A	ND	N/An
Heavier than TBA	0.005	N/A	ND	N/An

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N/An is not analyzed

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Tri-Ethylene Glycol Extra	ction Solvent:			
Component	Feed wt%	Solvent wt%	Overhead wt%	Bottoms wt%
Water	0.1800	0.0640	0.0190	1.0264
(WATER FREE BASIS)				
Lighter than MeOH/MF	0.207	ND	0.005	N/An
MeOH/MF	0.218	ND	0.004	N/An
PO	98.708	ND	99.991	N/An
Propanol/Propenal	ND	ND	ND	N/An
Acetone	0.754	ND	ND	N/An
TBA	0.112	ND	ND	N/An
Heavier than TBA	0.001	1.063	ND	N/An
Triethylene Glycol	ND	98.937	ND	N/An

Note: ND is not detected N/An is not analyzed

Solvent:Feed ratio 0.697 (kg/kg)

These distillations demonstrate that the triethylene glycol solvent improves the separation of methanol and water from propylene oxide.

Example II

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A glass 5.08cm (2") diameter vacuum jacketed Aldushan extractive distillation column having 120 actual plates was used for additional runs. The crude propylene oxide feed of Example I was introduced on tray 80 (from the top) and the triethylene glycol extractive distillation solvent was introduced on tray 30. The solvent:feed ratio was 5:1 by weight. The tower operating pressure was 159kPa (23 psia) at the condenser. Results show essentially complete removal of water and acetone from the feed propylene oxide. In addition, the methanol is substantially reduced in the overhead propylene oxide product, as illustrated by the following tables:

45 50	40		<i>30</i> <i>35</i>	25	20	15	10	5
				TABLE I				
		EXTRA	CTIVE DIST	EXTRACTIVE DISTILLATION OF IMPURE PROPYLENE OXIDE WITH TRIETHYLENE GLYCOL	MPURE PROPYLIE GLYCOL	ENE OXIDE		
			Over	Overheads Compositions.	tions, wt.			
Run: Number:		0	ო	4	വ	9	7	, co
Propylene Oxide (wt%)	99.860	99.890	906.66	99.619	99.877	99,636	99.507	99.621
Methanol (ppm)		434.888	400.273	412.137	550.98	591.317	663.873	485.332
Acetone (ppm)		6.003	2.412	41.406	4.132	8.287		
Water (ppm)	200	125	140	100	170	96	110	95
						-		
Run Number	თ	10	11	12	13	14	15	16
Propylene Oxide (wt%)	99.654		99,659		99.673	99,358	99.610	99.498
Methanol(ppm)) 567.		553:554	354.169	581.637	539.877	778.598	480.445
Acetone (ppm)	1.064		0.688	1.711	0.733		18.665	
Water (ppm)	120	110	06	110	94			

No bottoms measurements taken on Runs 4, 12 , 1	taken on	urements	toms meas	No bot	e glycol	s triethyler	* TEG represents triethylene glycol
86.204	86.030	86.381	84.248	83.324	85.231	84.548	TEG* (wt%)
				7800	7300	0069	Water (ppm)
			4.281	4.495	3,809	3.991	Acetone (ppm)
0.964	1.299	1.325	1.343	1.446	1,253	1.226	Methanol (ppm)
4.979	4.473	3.876	5.350	5.684	5.032	5.292	Propylene Oxide (wt%)
16	15	14	11	10	σ	æ	Run
84.487	84.692	85.948		85.786			TEG* (wt%)
		4800		3400	2500	1360	Water (ppm)
3.713	2.867	2,533		1.884			Acetone (ppm)
1.161	0.928	0.808		0.650			Methanol (ppm)
5.396	5.862	5.209		6.751			Propylene Oxide (wt%)
7	9	Ŋ		ю	. 73	.	Run <u>Number</u>
		-at	tions, wt.	Bottome Compositions, wt. &	ă.		
	30	PYLENE OXIC	IMPURE PRO	EXTRACTIVE DISTILLATION OF IMPURE PROPYLENE OXIDE WITH TRIETHYLENE GLYCOL	TRACTIVE DI	×	
			Ħ	TABLE II			
5	10	15	20	25 30	35	40	45 50

Claims

55 1. An extractive distillation process for the removal of oxygenated contaminants, including water, methanol, and acetone from an impure propylene oxide feedstock comprising propylene oxide contaminated with water, methanol and acetone, characterised by introducing the said impure propylene oxide feedstock into the lower half of a distillation column, introducing an extractive distillation agent

comprising triethylene glycol above the impure propylene oxide feed point, at a ratio of feedstock to extractive distillation agent of from 1:1 to 20:1, withdrawing from said distillation column an overhead distillate fraction comprising essentially anhydrous propylene oxide contaminated with reduced quantities of acetone and methanol, and a bottoms distillation fraction containing substantially all of the triethylene glycol, water and acetone and a portion of the methanol introduced into said distillation column.

- 2. A process as claimed in Claim 1 wherein the impure propylene oxide is charged to the lower half of a distillation column containing at least 25 theoretical plates and the extractive distillation agent is added at a point at least 4 theoretical plates above the impure propylene oxide charge point.
- 3. A process as claimed in Claim 1 or Claim 2 wherein the distillation conditions comprise a pressure of 69 to 276 kPa (10 to 40 psia), a reflux ratio of from 1:1 to 5:1, and a reboiler temperature within the range of 100 ° to 250 ° C. and a top temperature of 20 ° to 80 ° C.

Patentansprüche

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- 1. Ein Extraktivdestillationsverfahren zur Entfernung von sauerstoffhaltigen Verunreinigungen, einschließlich Wasser, Methanol und Aceton, aus einer unreinen Propylenoxid-Charge, die Propylenoxid umfaßt, das mit Wasser, Methanol und Aceton verunreinigt ist, dadurch gekennzeichnet, daß die besagte unreine Propylenoxid-Charge in die untere Hälfte einer Destillationskolonne eingeführt wird, ein Extraktionsmittel, das Triethylenglykol umfaßt, oberhalb des Zuführpunktes des unreinen Propylenoxids eingeführt wird, in einem Verhältnis von Charge zu Extraktionsmittel von 1:1 bis 20:1, aus besagter Destillationskolonne eine Überkopf-Destillatfraktion abgezogen wird, die im wesentlichen wasserfreies Propylenoxid umfaßt, das mit verringerten Mengen an Aceton und Methanol verunreinigt ist, und eine Bodenkörper-Destillationsfraktion, die im wesentlichen das gesamte Triethylenglykol, Wasser und Aceton und einen Teil des in besagte Destillationskolonne eingeführten Methanols enthält.
- 2. Ein Verfahren nach Anspruch 1, wobei das unreine Propylenoxid der unteren Hälfte einer Destillationskolonne zugeführt wird, die wenigstens 25 theoretische Böden enthält, und das Extraktionsmittel an einem Punkt wenigstens 4 theoretische Böden oberhalb des Zuführpunktes des unreinen Propylenoxids zugegeben wird.
- 3. Ein Verfahren nach Anspruch 1 oder Anspruch 2, wobei die Destillationsbedingungen einen Druck von 69 bis 276 kPa (10 bis 40 psia), ein Rückflußverhältnis von 1:1 bis 5:1 und eine Blasentemperatur im Bereich von 100 * bis 250 *C und eine Kopftemperatur von 20 * bis 80 *C umfassen.

Revendications

- 40 1. Procédé de distillation extractive pour l'élimination d'impuretés oxygénées, y compris l'eau, le méthanol et l'acétone d'une charge d'oxyde de propylène impure comprenant de l'oxyde de propylène contaminé par de l'eau, du méthanol et de l'acétone, caractérisé en ce qu'on introduit ladite charge d'oxyde de propylène impure dans la moitié inférieure de la colonne de distillation, en ce qu'on introduit un agent de distillation extractive comprenant du triéthylène glycol au-dessus du point de charge de l'oxyde de propylène impur, dans un rapport de la charge à l'agent de distillation extractive compris entre 1:1 et 20:1, en ce qu'on retire de ladite tête de distillation une fraction de distillat de tête comprenant essentiellement de l'oxyde de propylène anhydre contaminé avec des quantités réduites d'acétone et de méthanol, et une fraction de distillation de queues contenant sensiblement la totalité du triéthylène glycol, de l'eau et de l'acétone et une partie du méthanol introduit dans ladite colonne de distillation.
 - 2. Procédé selon la revendication 1 dans lequel l'oxyde de propylène impur est chargé dans la moitié inférieure d'une colonne de distillation contenant au moins 25 plateaux théoriques, et l'agent de distillation extractive est ajouté en un point situé à au moins 4 plateaux théoriques du point de charge de l'oxyde de propylène impur.
 - 3. Procédé selon la revendication 1 ou la revendication 2 dans lequel les conditions de distillation comprennent une pression de 69 à 276 kPa (10 à 40 psia), un rapport de reflux compris entre 1:1 et 5:1, et une température de rebouilleur comprise entre 100° et 250°C et une température au point

supérieur comprise entre 20 et 80 ° C.

